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Description**Field of the Invention**

5 This invention relates to peptides possessing selective ribonucleotide reductase inhibiting properties, to processes for their production, to pharmaceutical compositions of the peptides, and to the use of the peptides to inhibit ribonucleotide reductase.

Background of the Invention

10 Ribonucleotide reductase (RR) is the enzyme responsible for the reductive conversion of ribonucleotides to deoxyribonucleotides. The latter conversion is the rate controlling step in the biosynthesis of deoxyribonucleic acid (DNA), an essential principle for cell replication. RR activity has been linked directly to the proliferation of normal and neoplastic cells, with significantly higher levels of RR activity 15 being found in neoplastic cells (see E. Takeda and G. Weber, Life Sciences, 28, 1007 (1981) and G. Weber et al., Adv. Enz. Reg., 19, 87 (1981). Hence, the inhibition of RR activity is a valid target in the search for agents which will prevent or ameliorate abnormal cell proliferation as occurs, for example, in neoplasia and psoriasis.

20 Several inhibitors of mammalian RR have been investigated as potential antineoplastic agents or antitumor agents; for example, see B. van't Riet et al., J. Med. Chem., 22, 589 (1979), J. G. Cory et al., Adv. Enz. Reg., 19, 139 (1981), and B. van't Riet et al., U.S. patent 4,448,730, issued May 15, 1984. However, none have proved to be entirely satisfactory in clinical trials and only one RR inhibitor, hydroxyurea, is available to the physician for use as an antineoplastic agent. The latter drug, nevertheless, finds limited use because of side-effects and because frequent and large doses are required to maintain an effective 25 concentration of the drug in vivo (see Van't Riet et al., J. Med. Chem., supra). Accordingly, there is a need for an effective and safe inhibitor of mammalian RR.

The present application discloses a new group of peptides which are potent inhibitors of mammalian RR. This attribute, together with a relative lack of toxicity, renders the peptides useful as agents for combatting disease states associated with abnormal cell proliferation.

30 Peptides have previously been reported to be inhibitors of RR, see for instance J.H. Subak-Sharpe et al., UK patent application 2185024, published July 8, 1987, E.A. Cohen et al., European patent application 246630, published November 25, 1987, and R. Freidinger et al., European patent application 292255, published November 23, 1988. However, unlike the present peptides, the previously reported peptides are inhibitors of a viral RR and not mammalian RR.

Summary of the Invention

The peptides of this invention are represented by formula 1

40 $Y-R^1-R^2-R^3-R^4-R^5-R^6-Z$ 1

wherein

- R¹ is Thr, Thr(OBzl) or N-Me-Val,
- R² is Leu, N-Me-Leu or Phe,
- 45 R³ is Asp, D-Asp, Asp(NMe₂), Asn, Gln, Gly or Phe,
- R⁴ is Ala, D-Ala, Val or Glu,
- R⁵ is Asp, D-Asp or N-Me-Asp,
- R⁶ is Phe or homoPhe,
- Y is Phe,
- 50 AcPhe,
- H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe,
- H-Asn-Ser-Thr-Glu-Asn-Ser-Phe,
- H-Thr-Glu-Asn-Ser-Phe,
- AcThr-Glu-Asn-Ser-Phe,
- 55 AcAsn-Ser-Phe,
- H-Pro-Thr-Glu-Asn-Ser-Phe,
- H-Ser-Pro-Thr-Glu-Asn-Ser-Phe, or
- H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe,

and Z is hydroxy; or a therapeutically acceptable salt thereof.

A preferred group of peptides is represented by formula 1 wherein

- Y is AcPhe,
- 5 R₁ is Thr, Thr(OBzl) or N-Me-Val,
- R₂ is Leu or N-Me-Leu
- R₃ is Asp or Asn,
- R₄ is Ala, Val or Glu,
- R₅ is Asp or N-Me-Asp, and
- 10 R₆ is Phe.

Included within the scope of this invention is a pharmaceutical composition for treating abnormal cell proliferation in a mammal, comprising a peptide of formula 1, or a therapeutically acceptable salt thereof, and a pharmaceutically or veterinarianly acceptable carrier.

Within the scope of this invention a method is included for preventing or ameliorating abnormal cell proliferation in a mammal which comprises administering to the mammal an effective amount of a peptide of formula 1, or a therapeutically acceptable salt thereof.

Also included within the scope of the invention is a method of inhibiting ribonucleotide reductase which comprises administering to a mammal carrying a tumor having a relatively high ribonucleotide reductase level of activity, an amount of a peptide of formula 1, or a therapeutically acceptable salt thereof, effective to inhibit ribonucleotide reductase.

Processes for preparing the peptides of formula 1 are described hereinafter.

Details of the Invention

25 GENERAL

The term "residue" with reference to an amino acid means a radical derived from the corresponding α -amino acid by eliminating the hydroxyl of the carboxy group and one hydrogen of the α -amino group.

In general, the abbreviations used herein for designating the amino acids and the protective groups are based on recommendations of the IUPAC-IUB Commission of Biochemical Nomenclature, see European Journal of Biochemistry, 138, 9 (1984). For instance, Met, Met(O), Val, Thr, Glu, Gln, Ala, Ile, Asp, Phe, Ser, Leu, Asn and Tyr represent the residues of L-methionine, L-methionine sulfoxide, L-valine, L-threonine, L-glutamic acid, L-glutamine, L-alanine, L-isoleucine, L-aspartic acid, L-phenylalanine, L-serine, L-leucine, L-asparagine and L-tyrosine, respectively.

35 The symbol "Ac", when used herein as a prefix to a three letter symbol for an amino acid residue, denotes the N-acetyl derivative of the amino acid; for example, "AcPhe" represents the residue of N-acetyl-L-phenylalanine. Likewise, the symbol "N-Me", when used herein as a prefix to a three letter symbol for an amino acid residue, denotes the N-methyl derivative of the amino acid; for example, N-Me-Val represents the residue of N-methyl-L-valine.

40 Other symbols used herein are:

- Thr(OBzl) for the residue of O³-benzyl-L-serine
- Asp(NMe₂) for the residue of N⁴, N⁴-dimethyl L-asparagine
- HomoPhe for the residue of L-homophenylalanine, i.e. 2(S)-amino-4-phenylbutanoic acid

The amino acid residues, of which the designation therefor is not preceded by "D-", possess the L-configuration, including those with prefixes such as lower alkanoyl and acetyl. The amino acid residues of which the designation is preceded by "D-" possess the D-configuration. The starting materials for providing the amino acid residues, usually the corresponding N^a-protected amino acids, are commercially available or can be prepared by conventional methods.

50 The term "pharmaceutically acceptable carrier" as used herein means a non-toxic, generally inert vehicle for the active ingredient, which does not adversely affect the ingredient.

The term "veterinarily acceptable carrier" as used herein means a physiologically acceptable vehicle for administering drug substances to domestic animals comprising one or more non-toxic pharmaceutically acceptable excipients which do not react with the drug substance or reduce its effectiveness.

55 The term "coupling agent" as used herein means an agent capable of effecting the dehydrative coupling of an amino acid or peptide free carboxy group with a free amino group of another amino acid or peptide to form an amide bond between the reactants. The agents promote or facilitate the dehydrative coupling by activating the carboxy group. Descriptions of such coupling agents and activated groups are included in general textbooks of peptide chemistry; for instance, E. Schröder and K.L. Lübke, "The

Peptides", Vol. 1, Academic Press, New York, N.Y., 1965, pp 2-128, and K.D. Kopple, "Peptides and Amino acids", W.A. Benjamin, Inc., New York, N.Y., 1966, pp 33-51. Examples of coupling agents are thionyl chloride, diphenylphosphoryl azide, dicyclohexylcarbodiimide, N-hydroxysuccinimide, or 1-hydroxybenzotriazole in the presence of dicyclohexylcarbodiimide. A very practical and useful coupling agent is (benzotriazol-1-yloxy)tris(dimethylamino)phosphonium hexafluorophosphate, described by B. Castro et al., Tetrahedron Letters, 1219(1975), see also D. Hudson, J. Org. Chem., 53, 617 (1988), either by itself or in the presence of 1-hydroxybenzotriazole.

PROCESS

10

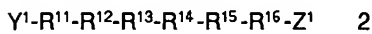
The peptides of formula 1 can be prepared by processes which incorporate therein methods commonly used in peptide synthesis such as classical solution coupling of amino acid residues and/or peptide fragments, and if desired solid phase techniques. Such methods are described, for example, by E. Schröder and K. Lübke, cited above, in the textbook series, "The Peptides: Analysis, Synthesis, Biology", 15 E. Gross et al., Eds., Academic Press, New York, N.Y., 1979-1987, Volumes 1 to 8, and by J.M. Stewart and J.D. Young in "Solid Phase Peptide Synthesis", 2nd ed., Pierce Chem. Co., Rockford, IL, USA, 1984.

15

A common feature of the aforementioned processes for the peptides is the protection of the labile side chain groups of the various amino acid residues with suitable protective groups which will prevent a chemical reaction from occurring at that site until the protective group is ultimately removed. Usually also 20 common feature is the protection of an α -amino group on an amino acid or a fragment while that entity reacts at the carboxy group, followed by the selective removal of the α -amino protective group to allow subsequent reaction to take place at that location. Usually another common feature is the initial protection of the C-terminal carboxyl of the amino acid residue or peptide fragment, which is to become the C-terminal function of the peptide, with a suitable protective group which will prevent a chemical reaction from 25 occurring at that site until the protective group is removed after the desired sequence of the peptide has been assembled.

25

Hence, the peptides of formula 1 can be prepared by a process comprising the stepwise coupling, in the order of the amino sequence of the peptide, of the appropriate amino acid residues or peptide fragments (with side chain functional groups duly protected, and with the C-terminal carboxyl of the amino 30 acid residue or peptide fragment, which is to become the C-terminal function of the peptide, duly protected by a C-terminal carboxyl protecting group), in the presence of a coupling agent, to obtain the protected peptide of formula 2



35

wherein R¹¹ is Thr(V¹) or N-Me-Val wherein V¹ is a protective group for the hydroxyl of Thr, R¹² has the same meaning as defined hereinabove for R², R¹³ is Asp(V²), D-Asp(V²), Asp(NMe₂), Asn, Gln, Gly or Phe wherein V² is a protective group for the ω -carboxyl of the amino acid residue designated therewith, R¹⁴ is Ala, D-Ala, Val or Glu(V²) wherein V² is as defined hereinabove, R¹⁵ is Asp(V²), D-Asp(V²) or N-Me-Asp(V²) 40 wherein V² is as defined hereinabove, R¹⁶ has the same meaning as defined hereinabove for R⁶, Y¹ is U-Phe or AcPhe wherein U is an α -amino protective group, or Y¹ is selected from the group of:

W¹-Val-Ile-Ser(V¹)-Asn-Ser(V¹)-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

W¹-Asn-Ser(V¹)-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

W¹-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

45

AcThr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

AcAsn-Ser(V¹)-Phe,

W¹-Pro-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

W¹-Ser(V¹)-Pro-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe or

50

W¹-Ser(V¹)-Ser(V¹)-Pro-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe, wherein V¹ is a protective group for the hydroxyl group of Thr or Ser, V² is a protective group for the ω -carboxyl of Glu and W¹ is an α -amino protective group or acetyl, and Z¹ is a classical carboxyl protective group or a resin support; followed by deprotecting (including cleaving the resin support if present), and acylating and/or amidating if required, the protected peptide of formula 2 to obtain the corresponding peptide of formula 1; and if desired, transforming the peptide of formula 1 into a therapeutically acceptable salt.

55

The term "resin support", as used herein with reference to Y¹, means the radical derived from a solid resin support of the type used in solid phase peptide synthesis. Such resin supports include the well known chloromethylated resins and benzhydrylamine resins, as well as resins which provide a spacer unit between the resin and the first amino acid building block of a peptide-resin system so that after the peptide portion

is assembled the resin can be cleaved selectively from the system. Examples of resins with spacers incorporated therein are α -(phenylacetamido)benzyl resin (PAB resin), described by E. Giralt et al., Tetrahedron 37, 2007 (1981), and 4-(2-bromo- or 4-(2-chloropropionyl)phenoxyacetyl BHA resins, photolabile resins described by D. Bellof and M. Mutter, Chemia, 39, 317 (1985).

5 Examples of side chain protective groups are benzyl for the protective group (V^1) for the hydroxyl of Thr or Ser; benzyl, 2,6-dichlorobenzyl or preferably cyclohexyl for the protective group (V^2) for the ω -carboxy of Asp or Glu and their related derivatives, and benzyl or preferably 2,6-dichlorobenzyl for the protective group (V^3) for the hydroxyl or Tyr. Note that when R^{11} of the protected peptide of formula 2 is Thr(OBzl) the benzyl group can serve a dual role, i.e. serve as the progenitor for the corresponding radical
10 in the ultimate product of formula 1 wherein R^1 is Thr(OBzl) or serve as a protective group. When the benzyl group is used as a progenitor, protective groups of the protected peptide of formula 2, if present, are those which can be removed selectively in the presence of benzyl by known methods.

Examples of C-terminal carboxyl protecting group include the classical groups, for example , benzyloxy and 4-nitrophenoxy, and for the present processes include also a "resin support".

15 In an embodiment of the exclusively solid phase method, the preparation of a peptide of formula 1 in which Z is hydroxy is commenced by coupling the first amino acid relative to the carboxy terminus (the amino acid having an α -amino protective group and, if required, a side chain protective group) with PAB resin in the presence of potassium fluoride or cesium chloride to give the corresponding solid resin support having the first amino acid (in protected form) linked thereto. The next step is the removal of the α -amino
20 protective group of the incorporated amino acid to give the free α -amino group. In the instance where the α -amino protective group is a t-butyloxycarbonyl, trifluoroacetic acid in methylene chloride or chloroform, or hydrochloric acid in dioxane, is used to effect the deprotection. The deprotection is carried out at a temperature between about 0 °C and room temperature. Other standard cleaving reagents and conditions for removal of specific α -amino protective groups may be used as described by E. Schröder and K. Lübke,
25 in "The Peptides", Vol. 1, Academic Press, New York, 1965, pp. 72-75. After removal of the α -amino protective group from the last mentioned intermediate, the remaining α -amino protected amino acids (with side chain protection when required) are coupled stepwise in the desired order to obtain the corresponding protected peptide of formula 2 attached to the PAB resin. Each protected amino acid is introduced into the reaction system in one to four fold excess and the coupling is effected with a coupling agent (one to three
30 fold excess) in a medium of methylene chloride, dimethylformamide, or mixtures of dimethylformamide and methylene chloride. In cases where incomplete coupling has occurred, the coupling procedure is repeated before removal of the α -amino protective group, prior to the coupling of the next protected amino acid. The success of the coupling reaction at each stage of the synthesis is monitored by the ninhydrin reaction as described by E. Kaiser et al., Anal. Biochem., 34, 595 (1970).

35 The preceding protected peptide of formula 2 thereafter is simultaneously cleaved from the resin and deprotected by treatment with liquid hydrogen fluoride to give the corresponding peptide of formula 1 in which Z is hydroxy.

When it is desired to prepare the C-terminal primary amide of formula 1 (Z = NH₂), the peptide can be prepared by the solid phase method using a benzhydrylamine resin and incorporating into the process the
40 cleavage of the resulting resin-bound peptide and any required deprotection according to known procedures such as described by Stewart and Young, supra.

Alternatively, a convenient and practical method for preparing the preceding C-terminal primary amide, as well as the corresponding secondary and tertiary amides (i.e. peptides of formula 1 wherein Z is lower alkylamino or di(lower alkyl)amino, respectively), involves the solid phase method with a photolabile resin
45 serving as the resin support. For instance, the stepwise coupling of the appropriate amino acid residues to 4-(2-chloropropionyl)phenoxyacetyl BHA resin, noted above, gives the protected peptide of formula 2 in which Z¹ is 4-(2-oxopropionyl)phenoxyacetyl BHA-resin. Subsequent photolysis of a suspension or solution of the latter peptide-resin (350 nm, 0 °C, 6 to 24 hours) gives the corresponding protected peptide of formula 2 in which Z¹ is hydroxy. Coupling of the latter protected peptide with benzylamine or the
50 appropriate lower alkylamine, e.g. methylamine or ethylamine, or the appropriate di(lower alkyl)amine, e.g. dimethylamine or ethylmethylamine, yields the respective protected peptide of formula 2 in which Z¹ is benzylamino, lower alkylamino or di(lower alkyl)amino. Deprotection of the latter protected peptide, for example with hydrofluoric acid, provides the corresponding C-terminal primary, secondary or tertiary amide of formula 1.

55 The terminal amino acylated derivatives of the peptides of formula 1, e.g. peptides of formula 1 wherein Y is (lower alkanoyl)-Phe or (lower alkanoyl)-Tyr, or Y is the decapeptide radical or fragment thereof wherein W is lower alkanoyl, are obtained from the corresponding free N-terminal amino peptide by treatment with a suitable acylating agent; for instance, the appropriate acid chloride or acid anhydride in the presence of a

strong organic base, e.g. 1-oxobutylchloride with diisopropylethylamine or N-methylmorpholine. Alternatively, the terminal amino acylated derivatives are obtained by using the appropriate N^a-acylated amino acid residue while preparing the peptide by conventional means. Again alternatively, the terminal amino acylated derivatives are obtained by coupling the corresponding free N-terminal amino peptide-resin (with side chain protection) with a molar equivalent of the appropriate lower alkanoic acid in the presence of a coupling agent; preferably (benzotriazol-1-yloxy)tris-(dimethylamino)phosphonium hexafluorophosphate, alone or in combination with 1-hydroxybenzotriazole, followed by conventional deprotection.

The peptide of formula 1 of this invention can be obtained in the form of therapeutically acceptable salts.

10 In the instance where a particular peptide has a residue which functions as a base, examples of such salts are those with organic acids, e.g. acetic, lactic, succinic, benzoic, salicylic, methanesulfonic or p-toluenesulfonic acid, as well as a polymeric acids such as tannic acid or carboxymethyl cellulose, and also salts with inorganic acids such as hydrohalic acid, or sulfuric acid, or phosphoric acid. If desired, a particular acid addition salt is converted into another acid addition salt, such as a non-toxic, pharmaceutically acceptable salt, by treatment with the appropriate ion exchange resin in the manner described by R.A. Boissonnas et al., *Helv. Chim. Acta*, 43, 1849 (1960).

In the instance where a particular peptide has one or more free carboxy groups, example of such salts are those with the sodium, potassium or calcium cations, or with strong organic bases, for example triethylamine or N-methylmorpholine.

20 In general, the therapeutically acceptable salts of the peptides of formula 1 are biologically fully equivalent to the peptides themselves.

BIOLOGICAL ASPECTS

25 The RR inhibiting and antineoplastic properties of the peptides of formula 1, or a therapeutically acceptable salt thereof, can be demonstrated by biochemical and biological procedures; for example, see H.L. Elford et al., *Adv. Enz. Reg.*, 19, 151 (1981). In the examples hereinafter, the RR inhibitory effect of exemplified peptides of formula 1 on human RR is demonstrated in the "Inhibition of Human Ribonucleotide Reductase Assay", the procedure of which is based on similar assays reported by E.A. Cohen, *J. Gen. Virol.*, 66, 733 (1985) and by Elford et al., *supra*.

Noteworthy is the finding that when the latter assay is repeated with other mammalian RR's and with RR's from bacterial and viral sources, a selective inhibition of mammalian RR is shown.

30 The ability of the peptides of formula 1 to selectively inhibit mammalian RR renders the peptides useful as agents for treating abnormal cell proliferation which occurs, for instance, in tumors (including both benign and malignant) and in psoriasis.

35 In the laboratory, the antineoplastic effect of the peptides can be demonstrated in tests with rodents having transplanted tumors. Survival time or tumor cell growth is used as the evaluation parameter. Examples of such transplantable tumors are lymphocytic leukemia, colon, mammary, melanocarcinoma and ependymoblastoma. The methods are described in various publications; for example, R.I. Geran et al, *Cancer Chemotherapy Report*, Part 3, 3, 1-103 (1972) and references therein.

40 When the peptides of this invention, or their therapeutically acceptable salts, are employed as agents for combatting disease states associated with abnormal cell proliferation, they are administered topically or systemically to warm-blooded animals, e.g. humans, dogs, horses, in combination with pharmaceutical acceptable carriers, the proportion of which is determined by the solubility and chemical nature of the peptide, chosen route of administration and standard biological practice. For example, for the treatment of psoriasis the peptide of formula 1 can be employed topically. For topical application, the peptides may be formulated in the form of solutions, creams, or lotions in pharmaceutically acceptable vehicles containing 1.0 - 10 per cent, preferably 2 to 5 per cent of the agent, and may be administered topically to the infected area of the body.

45 50 For systemic administration, the peptides of formula 1 are administered by either intravenous, subcutaneous or intramuscular injection, in compositions with pharmaceutically acceptable vehicles or carriers. For administration by injection, it is preferred to use the peptides in solution in a sterile aqueous vehicle which may also contain other solutes such as buffer or preservatives as well as sufficient quantities of pharmaceutically acceptable salts or of glucose to make the solution isotonic.

55 Examples of suitable excipients or carriers are found in standards pharmaceutical texts, e.g. in "Remington's Pharmaceutical Sciences", 16th ed, Mack Publishing Company, Easton, Penn., USA, 1980.

The dosage of the peptides will vary with the form of administration and the particular compound chosen. Furthermore, it will vary with the particular host under treatment. Generally, treatment is initiated

with small dosages substantially less than the optimum dose of the compound. Thereafter, the dosage is increased by small increments until the optimum effect under the circumstances is reached. In general, the peptides of this invention are most desirably administered at a concentration level that will generally afford effective results, without causing any harmful or deleterious side effects.

5 When used systemically as an antineoplastic or antitumor agent, the peptide of formula 1 is administered at a dose of 100 mcg to 1000 mcg per kilogram of body weight per day, although the aforementioned variations will occur. However, a dosage level that is in the range of from about 100 mcg to 500 mcg per kilogram of body weight per day is most desirably employed in order to achieve effective results.

10 The following examples illustrate further this invention. Solution percentages or ratios express volume to volume relationship, unless stated otherwise. Abbreviations used in the examples include Boc: t-butyloxycarbonyl; BOP: (benzotriazol-1-yloxy)tris(dimethylamino)phosphonium hexafluorophosphate; Bzl: benzyl; CH₂Cl₂: methylene chloride; Chxl: cyclohexyl; 2,6-DiClBzl: 2,6-dichlorobenzyl; DCC: N,N'-dicyclohexylcarbodiimide; DMF: dimethylformamide; HF: hydrofluoric acid; Et₂O: diethyl ether; EtOH: ethanol; HOBT: 1-hydroxy-benzotriazole; MeOH: methanol; TFA: trifluoroacetic acid.

Example 1

Preparation of Boc-Phe-CH₂-PAB resin

20 Boc-Phe-OH (29.7 g, 112 mmol) and potassium fluoride (15.7 g, 252 mmol) were added to a mechanically stirred suspension of α -(4-chloromethylphenylacetamido)benzyl copoly(styrene-1% divinylbenzene) resin (50 g, 28 mmol, described by Giralt et al., supra) in DMF (600 ml). The mixture was stirred at 70 °C for 24h, and then allowed to cool to ambient temperature. The solid was collected by filtration, 25 washed successively with 100 ml portions of DMF, DMF-H₂O(1:1), H₂O, H₂O-dioxane(1:1), dioxane, MeOH, CH₂Cl₂ and EtOH, and dried under reduced pressure to give 54.4 g of the title compound. The phenylalanine content of the product was 0.54 mmol/g as determined by deprotection of an aliquot and picric acid titration according to the method of B.F. Gisin, Anal. Chim. Acta, 58, 248 (1972).

30 Example 2

Preparation of the N-acetyl-heptapeptide of the formula:

AcPhe-Thr-Leu-Asp-Ala-Asp-Phe-OH

35 The title compound was synthetized by a modification of the solid-phase method of R.B. Merrifield, J. Amer. Chem. Soc., 85, 2149 (1963). Applying the method, the corresponding protected heptapeptide-resin having the correct sequence of amino acid residues was assembled by stepwise addition of the amino acids residues to Boc-Phe-CH₂-PAB resin, i.e. the title compound of Example 1. The following protocol was 40 used: (a) Boc-deprotection: 30% TFA in CH₂Cl₂ (2 times, firstly for 5 min then for 25 min); (b) wash: CH₂Cl₂ (3 times for 2 min each); (c) wash: isopropanol (2 min); (d) neutralization: 5% diisopropylethylamine in CH₂Cl₂ (2 times for 2 min each); (e) amino acid coupling: achieved by the method of D. Hudson, J. Org. Chem., 53, 617 (1988) using the appropriate protected amino acid (2.1 molar equivalents per mmol of the Boc-Phe-CH₂-PAB resin) and BOP-HOBT (2.2 and 1.1 molar equivalents, respectively, per mmol of the 45 Boc-Phe-CH₂-PAB resin) in the presence of N-methylmorpholine (6-8 molar equivalents providing pH 8 for the reaction mixture) in CH₂Cl₂ or DMF; the reaction time for coupling varied from 3 to 5h; and (f) wash: CH₂Cl₂ or DMF (2 times for 2 min each. The Gln and Asn residues were coupled in DMF after activation of the corresponding Boc-amino acid with DCC-HOBT and removal by filtration of the N,N'-dicyclohexylurea formed during the activation process.

50 The Boc group gave N^o protection for all amino acids. Side chain protection was as follows: Bzl for Thr and Ser, Chxl for Asp and Glu, and 2,6-DiClBzl for Tyr. After each coupling, the completeness of the reaction was checked by the ninhydrin test, E. Kaiser et al., Anal. Biochem., 34, 595 (1970). The N-terminal acetylation was accomplished by coupling the free N-terminal amino protected peptide-resin with a molar equivalent of acetic acid using the BOP-HOBT method, or with acetic anhydride in the presence of 55 diisopropylethylamine in CH₂Cl₂ or DMF.

On completion of the peptide sequence, the protected heptapeptide-resin was collected on a filter, washed with CH₂Cl₂ and EtOH and dried under reduced pressure over phosphorus pentoxide for 24 h to give the corresponding protected heptapeptide-resin (i.e. peptide-resin). The heptapeptide was cleaved from

the peptide-resin by using HF (5 ml per g of peptide-resin) in the presence of distilled anisole (1 ml per g of peptide-resin) and ethanedithiol (0.2 ml per g of peptide-resin). The mixture was maintained at -20°C for 40 min and then at 0-5°C for 40 min, with vigorous stirring. After evaporation of HF, the residue was triturated with Et₂O. The mixture was filtered through distomaceous earth (Celite®). After washing with Et₂O, the filter cake was dried under reduced pressure. The residual solid was washed with several portions of 10% aqueous acetic acid, and then with 0.1M aqueous NH₄OH (total volume: 40 ml per g of the peptide-resin). All the aqueous filtrates were mixed at 0°C (pH 6) and lyophilized to afford a white solid residue.

Purification of the solid residue to greater than 95% homogeneity was accomplished by reversed phase HPLC with a Waters model 600 multisolvent delivery system (Waters, Milford, MA, USA) equipped with a 10 UV detector and using a Whatman Partisil® 100DS-3 C-18 column (2.2 x 50 cm²), 10 micron particle size. The elution was done with a gradient of acetonitrile in 0.1% aqueous TFA such as:

- a) initial: 10% acetonitrile in 0.1% aqueous TFA for 20 min,
- b) followed by gradually increasing the concentration of acetonitrile to 20% over a period of 20 min, followed by gradually increasing the concentration to 40% acetonitrile over a period of 50 min.

Pure fractions, as determined by analytical HPLC, were pooled and lyophilized to afford the title heptapeptide as a trifluoroacetate salt. Analytical HPLC showed the product to be at least 95% homogeneous. Amino acid analysis: Phe, 2.00; Asp, 2.06; Thr, 0.95; Leu, 0.99; Ala, 1.00; FAB-MS, calcd: 869.38, found: 870 (M + H), 892 (M + Na), etc.

Example 3

Inhibition of Human Ribonucleotide Reductase Assay

1) Preparation of extracts containing active RR:

- (a) Cell line: HeLa cells (ATCC CCL 2.2), human epitheloid carcinoma, cervix.
- (b) Cell culture: Cells were incubated in a medium consisting of Iscove's Modified Dulbecco Medium, pH 7.2-7.4 (Gibco Canada Inc., Burlington, ON, Canada) supplemented with 10% by volume of fetal calf serum, heat inactivated (Gibco Canada Inc.), 1 millimolar (mM) of sodium pyruvate, 100 units/ml of penicillin G, 100 g/ml of streptomycin and 2 mM of L-glutamine. The incubation was done in a 1 l spinner flask at 37°C under a mixture of 5% CO₂ in air. Cell-containing media (75% of culture volume) was withdrawn semicontinuously at 24-48h intervals. Fresh media was added each time to replace withdrawn media. Final cell density at harvest was 1-2 x 10⁶ cells/ml.
- (c) Preparation of cell extract containing human ribonucleotide reductase (hRR):

The harvested culture media obtained above was subjected to low speed centrifugation. The resulting cell pellet was processed according to the following steps. (All steps were performed at 4°C unless noted otherwise.)

Step

- 1) Wash Buffer 100 mM KH₂PO₄/K₂HPO₄ in 0.9% (w/v) sodium chloride (pH 7.2).
- 2) Storage Cells frozen at -80°C until extraction.
- 3) Extraction Buffer 25 mM N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid (HEPES) buffer (pH 7.6), 2 mM DL-dithiothreitol (DTT) and 1 mM MgCl₂.
- 4) Cell Disruption Cells in extraction buffer held for 30 min at 4°C followed by 20 strokes on a Potter-Elvehjem Homogenizer (Kontes Glass Co., Vineland, NJ, USA).
- 5) Centrifugation 40,000 times gravity for 60 min; recover supernatant.
- 6) Precipitation A solution of 5% (w/v) streptomycin sulphate in 50 mM HEPES, 5 mM DTT and 5 mM MgCl₂ added dropwise to supernatant to give final concentration in mixture of 1% (w/v) of streptomycin sulfate.
- 7) Centrifugation 40,000 times gravity for 60 min; recover supernatant.
- 8) Precipitation Saturated (NH₄)₂SO₄ in HEPES/DTT/MgCl₂ buffer (see step 6) added slowly to supernatant to yield 50% saturated solution; solution agitated for 30 min.
- 9) Centrifugation 40,000 times gravity for 60 min.; recover pellet.
- 10) Solubilization Take up pellet in minimum volume of HEPES/DTT/MgCl₂ buffer (see step 6).
- 11) Dialysis 3-Cycle diafiltration (1h per cycle) carried out against HEPES/DTT/MgCl₂ buffer (see step 6) using a microconcentrator with a 10,000 MW cut-off (Centricon® 10, Amicon, Danvers, MA, USA).

12) Storage Frozen at -80 °C.

2) Assay Protocol:

5 (a) Standard Reaction Mixture:

	Component	Amount*
10	HEPES Buffer (pH 7.8)	50 mM
	Adenosine Triphosphate	4 mM
	DTT	30 mM
	MgCl ₂	11.5 mM
	NaF	4 mM
15	Cytidine Diphosphate (CDP)	0.054 mM
	¹⁴ C CDP (DuPont Chemical Co. Lachine, QC, Canada)	0.17 Ci/ml
	Bacitracin	1 mM
	Test Compound	1-250 M

* Final concentration of component in standard reaction mixture.

20

(b) Assay Procedure:

The activity of RR was quantitated by following the conversion of radiolabeled cytidine diphosphate to radiolabeled deoxycytidine diphosphate, i.e. (¹⁴C)CDP to (¹⁴C)dCDP. The amount of cell extract utilized in the assay was that which gave a linear response between enzyme concentration and CDP conversion (ca. 200 g of protein per assay).

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After addition of the cell extract, the assay mixture was incubated at 37 °C for 30 min. The reaction was stopped by immersing the vessel containing the assay mixture in boiling water for 4 min. Nucleotides in the supernatant were then converted to nucleosides by the addition of excess *Crotalus adamanteus* snake venom (ca. 20 l of a preparation of 40 mg/ml of the venom in an aqueous solution of 14 mM tris-(hydroxymethyl)aminomethane (pH 8.8) and 46.5 mM MgCl₂), followed by incubating the resulting mixture for 60 min at 37 °C. The enzymatic reaction was stopped by immersing the vessel containing the reaction mixture in boiling water for 6 min. Thereafter, the mixture is centrifuged at 10,000 rpm on a clinical centrifuge for 5 min.

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The resulting free nucleosides, cytidine (C) and deoxycytidine (dC), in the supernatant were separated by thin layer chromatography on polyethyleneimine-cellulose plates pretreated with boric acid. Elution of 5 l samples was accomplished using a solution of ethanol / 20 mM aqueous ammonium formate (1:1), pH 4.7. Quantitation of radiolabel migrating as C and dC was carried out using radioanalytical imaging equipment (AMBIS Systems Inc., San Diego, CA, USA). Substrate conversion was calculated as:

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(¹⁴C) deoxycytidine

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(¹⁴C) deoxycytidine + (¹⁴C) cytidine

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A unit of ribonucleotide reductase activity is defined as that amount which reduces one nmole of CDP/minute under the conditions described above. Activity was calculated from substrate conversion using the following relationship:

$$\begin{array}{c}
 5 \\
 \left[\begin{array}{l} \text{substrate} \\ \text{conversion} \\ (\text{sample}) \end{array} \right] - \left[\begin{array}{l} \text{substrate} \\ \text{conversion} \\ (\text{blank}) \end{array} \right] \times \text{conversion factor} = \\
 10 \\
 \text{activity units}
 \end{array}$$

15 The conversion factor for the Hela assay was 0.108. Specific activity was expressed as units/mg of protein in the incubation mixture. In one embodiment, the specific activity of the Hela extract was found to be 0.2 units/mg.

The peptides of formula 1 were tested at a minimum of three concentrations. IC₅₀'s were estimated from graphs plotting the results for each peptide, the IC₅₀ being the concentration of the peptide in micromoles (M) producing 50% of the maximal inhibition of the enzyme.

20 When the N-acetyl-heptapeptide of Example 2 having the formula AcPhe-Thr-Leu-Asp-Ala-Asp-Phe-OH was tested according to the assay of this example, an IC₅₀ of 38 M was determined for the compound.

Example 4

25 The following table of exemplified peptides of formula 1 further illustrates the invention. The peptides, prepared in a manner analogous to that described for the N-acetyl-heptapeptide of Example 2, are listed with their characterizing physical data and their IC₅₀ as determined by the inhibition of human RR assay of Example 3.

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	Peptide	Amino Acid Analysis	FAB/MS		IC_{50} (M)
			Calcd (FW)	Found	
5					
10	H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH	Asp+Asn, 4.17; Thr, 1.97; Ser, 2.59; Glu, 0.97; Ala, 1.06; Val, 0.42; Ile, 0.40; Leu, 1.1; Phe, 2.14		1758.8	1759 ¹
15					49
20	H-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH	Asp+Asn, 4.19; Thr, 1.94; Ser, 1.80; Glu, 1.0; Ala, 0.99; Leu, 1.01; Phe, 2.05		1459.6	1460 ¹ 1483 ²
25	H-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH	Asp+Asn, 3.08; Thr, 1.95; Ser, 0.92; Glu, 1.01; Ala, 1.0; Leu, 1.01; Phe, 2.03		1258.5	1259 ¹ 1281 ²
30	AcThr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH			1301 ¹ 1300.6	95 1323 ²
35	Ac-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH			1070.5	1093 ²
40					33

¹Protonated parent ion (M+1)

²Parent ion associated with Na⁺(M+23)

The capacity of the peptides of formula 1, noted in Example 1 and listed above, to inhibit the enzymatic action of hamster RR was demonstrated by a variation of the RR assay of Example 2 wherein the cell extract of human RR is replaced by an extract of hamster RR. The latter extract was prepared according to the procedure of Cohen et al., supra, from hamster 600H cells obtained from an overproducing strain of Chinese hamster lung cell line, selected for hydroxyurea resistance.

Other examples of peptides within the scope of this invention are listed hereinafter together with their IC_{50} (shown in parenthesis) as determined in the inhibition of human RR assay:

5 AcPhe-Thr(OBzl)-Leu-Asp-Ala-Asp-Phe-OH (28),
AcPhe-Thr-Leu-Asp-Ala-N-Me-Asp-Phe-OH (46),
AcPhe-Thr-Phe-Asn-Glu-Asp-Phe-OH (150),
AcPhe-Thr-Leu-Asp-Ala-D-Asp-Phe-OH (280),
10 AcPhe-Thr-N-Me-Leu-Asp-Ala-Asp-Phe-OH (23),
AcPhe-N-Me-Val-Leu-Asp-Ala-Asp-Phe-OH (48),
AcPhe-Thr-Leu-Asp-D-Ala-Asp-Phe-OH (460),
AcPhe-Thr-Leu-D-Asp-Ala-Asp-Phe-OH (200),
15 AcPhe-Thr-Leu-Asp-Val-Asp-Phe-OH (39),
AcPhe-Thr-Leu-Gln-Ala-Asp-Phe-OH (120),
AcPhe-Thr-Leu-Asn-Ala-Asp-Phe-OH (31),
20 AcPhe-Thr-Leu-Asp-Ala-Asp-homoPhe-OH (58),
AcPhe-Thr-Leu-Asp-Glu-Asp-Phe-OH (28),
AcPhe-Thr-Leu-Asp(NMe₂)-Ala-Asp-Phe-OH (58),
25 AcPhe-Thr-Leu-Gly-Ala-Asp-Phe-OH (150),
AcPhe-Thr-Leu-Phe-Ala-Asp-Phe-OH (73),
30 H-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH (41),
H-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH
(29), and
35 H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-
Ala-Asp-Phe-OH (26).

40 Claims

1. A compound of formula 1

Y-R¹-R²-R³-R⁴-R⁵-R⁶-Z 1

wherein

R¹ is Thr, Thr(OBzl) or N-Me-Val,

R² is Leu, N-Me-Leu or Phe,

R^3 is Asp, D-Asp, Asp(NMe₂), Asn, Gln, Gly or Phe,

R⁴ is Ala, D-Ala, Val or Glu,

R^5 is Asp, D-Asp or N-Me-Asp

R⁶ is Phe or homoPhe,

Y is Phe,

AcPhe,

H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe,

H-Asn-Ser-Thr-Glu-Asn-Ser-Phe,

H-Thr-Glu-Asn-Ser-Phe,

AcThr-Glu-Asn-Ser-Phe,

AcAsn-Ser-Phe,
H-Pro-Thr-Glu-Asn-Ser-Phe,
H-Ser-Pro-Thr-Glu-Asn-Ser-Phe, or
H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe,
5 and Z is hydroxy; or a therapeutically acceptable salt thereof.

2. A peptide as recited in claim 1 wherein

Y is AcPhe,
R₁ is Thr, Thr(OBzl) or N-Me-Val,
10 R₂ is Leu or N-Me-Leu
R₃ is Asp or Asn,
R₄ is Ala, Val or Glu,
R₅ is Asp or N-Me-Asp, and
R₆ is Phe.

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3. A peptide of formula 1 of claim 1 selected from the group of:

5 AcPhe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-
Asp-Ala-Asp-Phe-OH,
10 H-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-
Phe-OH,
H-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
15 AcThr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
AcAsn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
AcPhe-Thr(OBzl)-Leu-Asp-Ala-Asp-Phe-OH,
20 AcPhe-Thr-Leu-Asp-Ala-N-Me-Asp-Phe-OH,
AcPhe-Thr-Phe-Asn-Glu-Asp-Phe-OH,
AcPhe-Thr-Leu-Asp-Ala-D-Asp-Phe-OH,
25 AcPhe-Thr-N-Me-Leu-Asp-Ala-Asp-Phe-OH,
AcPhe-N-Me-Val-Leu-Asp-Ala-Asp-Phe-OH,
Ac-Phe-Thr-Leu-Asp-D-Ala-Asp-Phe-OH,
30 AcPhe-Thr-Leu-D-Asp-Ala-Asp-Phe-OH,
AcPhe-Thr-Leu-Asp-Val-Asp-Phe-OH,
AcPhe-Thr-Leu-Gln-Ala-Asp-Phe-OH,
35 AcPhe-Thr-Leu-Asn-Ala-Asp-Phe-OH,
AcPhe-Thr-Leu-Asp-Ala-Asp-homoPhe-OH,
AcPhe-Thr-Leu-Asp-Glu-Asp-Phe-OH,
40 AcPhe-Thr-Leu-Asp(NMe₂)-Ala-Asp-Phe-OH,
AcPhe-Thr-Leu-Gly-Ala-Asp-Phe-OH,
AcPhe-Thr-Leu-Phe-Ala-Asp-Phe-OH,
45 H-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
H-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-
Phe-OH,
50 H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-
Asp-Phe-OH.

55 4. A pharmaceutical composition comprising a peptide as recited in any of the claims 1 to 3, or a therapeutically acceptable salt thereof, and a pharmaceutically or veterinarily acceptable carrier.

5. Use of a peptide or composition as recited in any of the claims 1 to 4 for the manufacture of a medicament for application as an antineoplastic or antitumor agent.

6. Use of a peptide or composition as recited in any of claims 1 to 4 for the manufacture of a medicament for application as a mammalian ribonucleotide reductase inhibitor.

5
7. A process for preparing a peptide of formula 1 of claim 1, which comprises deprotecting the protected peptide of formula 2

10 $Y^1-R^{11}-R^{12}-R^{13}-R^{14}-R^{15}-R^{16}-Z^1$ 2

wherein R^{11} is Thr(V^1) or N-Me-Val wherein V^1 is a protective group for the hydroxyl of Thr, R^{12} has the same meaning as defined for R^2 in claim 1, R^{13} is Asp(V^2), D-Asp(V^2), Asp(NMe₂), Asn, Gln, Gly or Phe wherein V^2 is a protective group for the ω -carboxyl of the amino-acid residue designated therewith, R^{14} is Ala, D-Ala, Val or Glu(V^2) wherein V^2 is as defined herein, R^{15} is Asp(V^2), D-Asp(V^2) or N-Me-Asp(V^2) wherein V^2 is as defined herein, R^{16} has the same meaning as defined for R^6 in claim 1, Y^1 is U-Phe or AcPhe wherein U is an α -amino protective group, or Y^1 is selected from the group of:
 W¹-Val-Ile-Ser(V^1)-Asn-Ser(V^1)-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 20 W¹-Asn-Ser(V^1)-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 W¹-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 AcThr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 AcAsn-Ser(V^1)-Phe,
 W¹-Pro-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 25 W¹-Ser(V^1)-Pro-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe or
 W¹-Ser(V^1)-Ser(V^1)-Pro-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe, wherein V^1 is a protective group for the hydroxyl group of Thr or Ser, V^2 is a protective group for the ω -carboxyl of Glu and W^1 is an α -amino protective group or acetyl, and Z^1 is a classical carboxyl protective group or a resin support; followed by deprotecting (including cleaving the resin support if present), and acylating and/or amidating if required, the protected peptide of formula 2 to obtain the corresponding peptide of formula 1; and if desired, transforming the peptide of formula 1 into a therapeutically acceptable salt.

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8. A process for preparing the protected peptide of formula 2 of claim 7 in which Z^1 is a carboxyl protective group or a resin support, which comprises:
 35 stepwise coupling in the order of the amino acid sequence of the protected peptide of formula 2, the protected amino acid residues or peptide fragments in which:
 i) labile side chain groups of the residues or fragments are protected with suitable protective groups to prevent chemical reactions from occurring at that site until the protective group is ultimately removed after the completion of the stepwise coupling;
 40 ii) an α -amino group of a coupling reactant is protected by an α -amino protective group while the free carboxy group of that reactant couples with the free α -amino group of the second reactant; the α -amino protective group being one which can be selectively removed to allow the subsequent coupling step to take place at that α -amino group; and
 iii) The C-terminal carboxyl of the amino acid residue of the amino acid residue or peptide fragment, which is to become the C-terminal function of the protected peptide, is protected with a suitable protective group which will prevent chemical reaction occurring at that site until after the desired amino acid sequence for the protected peptide has been assembled;
 45 to obtain the protected peptide of formula 2.

50 Patentansprüche

1. Verbindung der Formel 1

55 $Y-R^1-R^2-R^3-R^4-R^5-R^6-Z$ 1

worin

R^1 Thr, Thr(OBzl) oder N-Me-Val ist,
 R^2 Leu, N-Me-Leu oder Phe ist,

R³ Asp, D-Asp, Asp(NMe₂), Asn, Gln, Gly oder Phe ist,
R⁴ Ala, D-Ala, Val oder Glu ist,
R⁵ Asp, D-Asp oder N-Me-Asp ist,
R⁶ Phe oder homoPhe ist,

5 Y Phe,

AcPhe

H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe,

H-Asn-Ser-Thr-Glu-Asn-Ser-Phe,

H-Thr-Glu-Asn-Ser-Phe,

10 AcThr-Glu-Asn-Ser-Phe,

AcAsn-Ser-Phe,

H-Pro-Thr-Glu-Asn-Ser-Phe,

H-Ser-Pro-Thr-Glu-Asn-Ser-Phe, oder

H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe ist,

15 und Z Hydroxy ist; oder ein therapeutisch annehmbares Salz dieser Verbindung.

2. Peptid nach Anspruch 1, worin

Y AcPhe ist,

R₁ Thr, Thr(OBzl) oder N-Me-Val ist,

20 R₂ Leu oder N-Me-Leu ist,

R₃ Asp oder Asn ist,

R₄ Ala, Val oder Glu ist,

R₅ Asp oder N-Me-Asp ist, und

R₆ Phe ist.

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3. Peptid der Formel 1 nach Anspruch 1, ausgewählt aus der Gruppe der folgenden Verbindungen:

AcPhe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 5 H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 Ala-Asp-Phe-OH,
 H-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 10 H-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 AcThr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 AcAsn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 AcPhe-Thr(OBzl)-Leu-Asp-Ala-Asp-Phe-OH,
 15 AcPhe-Thr-Leu-Asp-Ala-N-Me-Asp-Phe-OH,
 AcPhe-Thr-Phe-Asn-Glu-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp-Ala-D-Asp-Phe-OH,
 20 AcPhe-Thr-N-Me-Leu-Asp-Ala-Asp-Phe-OH,
 AcPhe-N-Me-Val-Leu-Asp-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp-D-Ala-Asp-Phe-OH,
 25 AcPhe-Thr-Leu-D-Asp-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp-Val-Asp-Phe-OH,
 AcPhe-Thr-Leu-Gln-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asn-Ala-Asp-Phe-OH,
 30 AcPhe-Thr-Leu-Asp-Ala-Asp-homoPhe-OH,
 AcPhe-Thr-Leu-Asp-Glu-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp(NMe₂)-Ala-Asp-Phe-OH,
 35 AcPhe-Thr-Leu-Gly-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Phe-Ala-Asp-Phe-OH,
 H-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 40 H-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,

H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH.

45 4. Pharmazeutische Zubereitung, enthaltend ein Peptid nach einem der Ansprüche 1 bis 3 oder ein therapeutisch annehmbares Salz desselben und einen pharmazeutisch oder veterinärmedizinisch annehmbaren Träger.

50 5. Verwendung eines Peptids oder einer Zubereitung nach einem der Ansprüche 1 bis 4 zur Herstellung eines Medikamentes für die Anwendung als antineoplastischer Wirkstoff oder als Antitumorwirkstoff.

6. Verwendung eines Peptids oder einer Zubereitung nach einem der Ansprüche 1 bis 4 zur Herstellung eines Medikamentes für die Anwendung als Säugetier-Ribonukleotidreduktase-Hemmer.

55 7. Verfahren zur Herstellung eines Peptids der Formel 1 nach Anspruch 1, welches die Entschützung des geschützten Peptids der Formel 2

$Y^1-R^{11}-R^{12}-R^{13}-R^{14}-R^{15}-R^{16}-Z^1$ 2

beinhaltet, worin R^{11} Thr(V^1) oder N-Me-Val ist, worin V^1 eine Schutzgruppe für das Hydroxyl von Thr ist, R^{12} die gleiche Bedeutung hat, wie für R^2 in Anspruch 1 definiert wurde, R^{13} Asp(V^2), D-Asp(V^2), Asp(NMe₂), Asn, Gln, Gly oder Phe ist, worin V^2 eine Schutzgruppe für das ω -Carboxyl des damit bezeichneten Aminosäurerestes ist, R^{14} Ala, D-Ala, Val oder Glu(V^2) ist, worin V^2 wie hier oben definiert ist, R^{15} Asp(V^2), D-Asp(V^2) oder N-Me-Asp(V^2) ist, worin V^2 wie hier oben definiert ist, R^{16} die gleiche Bedeutung hat, wie für R^6 in Anspruch 1 definiert wurde, Y^1 U-Phe oder AcPhe ist, worin U eine α -Amino-Schutzgruppe ist, oder Y^1 ausgewählt ist aus der Gruppe der folgenden Verbindungen:

10 W^1 -Val-Ile-Ser(V^1)-Asn-Ser(V^1)-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 W^1 -Asn-Ser(V^1)-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 W^1 -Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
AcThr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
AcAsn-Ser(V^1)-Phe,

15 W^1 -Pro-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,
 W^1 -Ser(V^1)-Pro-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe oder
 W^1 -Ser(V^1)-Ser(V^1)-Pro-Thr(V^1)-Glu(V^2)-Asn-Ser(V^1)-Phe,

20 worin V^1 eine Schutzgruppe für die Hydroxylgruppe von Thr oder Ser ist, V^2 eine Schutzgruppe für das ω -Carboxyl von Glu ist und W^1 eine α -Amino-Schutzgruppe oder Acetyl ist, und Z^1 eine klassische Carboxyl-Schutzgruppe oder ein Harzträger ist; gefolgt von der Entschüttung (einschliesslich der Abspaltung des allenfalls vorhandenen Harzträgers) und erforderlichenfalls der Acylierung und/oder Amidierung des geschützten Peptids der Formel 2 zur Gewinnung des entsprechenden Peptids der Formel 1; und gewünschtenfalls der Umwandlung des Peptids der Formel 1 in ein therapeutisch annehmbares Salz.

25 8. Verfahren zur Herstellung des geschützten Peptids der Formel 2 von Anspruch 7, worin Z^1 eine Carboxyl-Schutzgruppe oder ein Harzträger ist, welches beinhaltet:
schriftweise Kupplung der geschützten Aminosäurereste oder Peptidfragmente gemäss der Reihenfolge der Aminosäuresequenz des geschützten Peptids der Formel 2, wobei:

30 i) labile Seitenkettengruppen der Reste oder Bruchstücke mit geeigneten Schutzgruppen geschützt werden, die das Auftreten von chemischen Reaktionen an dieser Stelle verhindern, bis die Schutzgruppe nach Abschluss der schrittweisen Kupplung endgültig entfernt wird;

35 ii) eine α -Aminogruppe eines Kupplungsmittels durch eine α -Amino-Schutzgruppe geschützt wird, während die freie Carboxygruppe dieses Mittels mit der freien α -Aminogruppe des zweiten Mittels gekuppelt wird; wobei die α -Amino-Schutzgruppe eine solche ist, die selektiv entfernt werden kann, damit der anschliessende Kupplungsschritt an dieser α -Aminogruppe stattfinden kann; und

40 iii) das C-terminale Carboxyl des Aminosäurerestes oder Peptidfragmentes, welches zur C-terminalen Funktion des geschützten Peptids werden soll, durch eine geeignete Schutzgruppe geschützt wird, die das Auftreten einer chemischen Reaktion an dieser Stelle verhindert, bis die für das geschützte Peptid gewünschte Aminosäuresequenz zusammengebaut wurde;
zur Gewinnung des geschützten Peptids der Formel 2.

Revendications

45 1. Composé de formule 1

$Y-R^1-R^2-R^3-R^4-R^5-R^6-Z$ 1

où

50 R^1 est Thr, Thr(OBzl) ou N-Me-Val,
 R^2 est Leu, N-Me-Leu ou Phe,
 R^3 est Asp, D-Asp, Asp(NMe₂), Asn, Gln, Gly ou Phe,
 R^4 est Ala, D-Ala, Val ou Glu,
 R^5 est Asp, D-Asp ou N-Me-Asp,
 R^6 est Phe ou homoPhe,
 Y est Phe,
AcPhe,
H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe,

H-Asn-Ser-Thr-Glu-Asn-Ser-Phe,
H-Thr-Glu-Asn-Ser-Phe,
AcThr-Glu-Asn-Ser-Phe,
AcAsn-Ser-Phe,
5 H-Pro-Thr-Glu-Asn-Ser-Phe,
H-Ser-Pro-Thr-Glu-Asn-Ser-Phe, ou
H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe,
et Z est hydroxyle; ou sel thérapeutiquement acceptable de celui-ci.

10 2. Peptide selon la revendication 1, dans lequel

Y est AcPhe,
R¹ est Thr, Thr(OBzl) ou N-Me-Val,
R² est Leu ou N-Me-Leu,
R³ est Asp ou Asn,
15 R⁴ est Ala, Val ou Glu,
R⁵ est Asp ou N-Me-Asp, et
R⁶ est Phe.

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3. Peptide de formule 1 selon la revendication 1 choisi dans le groupe suivant:

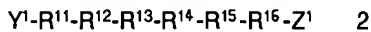
AcPhe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 5 H-Val-Ile-Ser-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 Ala-Asp-Phe-OH,
 10 H-Asn-Ser-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 H-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 15 AcThr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 AcAsn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 20 AcPhe-Thr(OBz1)-Leu-Asp-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp-Ala-N-Me-Asp-Phe-OH,
 AcPhe-Thr-Phe-Asn-Glu-Asp-Phe-OH,
 25 AcPhe-Thr-Leu-Asp-Ala-D-Asp-Phe-OH,
 AcPhe-Thr-N-Me-Leu-Asp-Ala-Asp-Phe-OH,
 AcPhe-N-Me-Val-Leu-Asp-Ala-Asp-Phe-OH,
 30 AcPhe-Thr-Leu-Asp-D-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-D-Asp-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp-Val-Asp-Phe-OH,
 35 AcPhe-Thr-Leu-Gln-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asn-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Asp-Ala-Asp-homoPhe-OH,
 AcPhe-Thr-Leu-Asp-Glu-Asp-Phe-OH,
 40 AcPhe-Thr-Leu-Asp(NMe₂)-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Gly-Ala-Asp-Phe-OH,
 AcPhe-Thr-Leu-Phe-Ala-Asp-Phe-OH,
 45 H-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 H-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH,
 H-Ser-Ser-Pro-Thr-Glu-Asn-Ser-Phe-Thr-Leu-Asp-Ala-Asp-Phe-OH.

4. Composition pharmaceutique comprenant un peptide selon l'une quelconque des revendications 1 à 3, ou un sel thérapeutiquement acceptable de celui-ci, et un véhicule acceptable du point de vue pharmaceutique ou vétérinaire.

5. Utilisation d'un peptide ou d'une composition selon l'une quelconque des revendications 1 à 4 pour la fabrication d'un médicament destiné à être appliqué comme agent antinéoplasique ou antitumoral.

6. Utilisation d'un peptide ou d'une composition selon l'une quelconque des revendications 1 à 4 pour la fabrication d'un médicament destiné à être appliqué comme inhibiteur de ribonucléotidé réductase de mammifère.

7. Procédé pour préparer un peptide de formule 1 selon la revendication 1, qui comprend la déprotection du peptide protégé de formule 2



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où R¹¹ est Thr(V¹) ou N-Me-Val où V¹ est un groupe protecteur pour l'hydroxyle de Thr, R¹² a la même signification que celle définie pour R² dans la revendication 1, R¹³ est Asp(V²), D-Asp(V²), Asp(NMe₂), Asn, Gln, Gly ou Phe où V² est un groupe protecteur pour le ω -carboxyle du résidu d'acide aminé représenté avec lui, R¹⁴ est Ala, D-Ala, Val ou Glu(V²) où V² est tel que défini ici, R¹⁵ est Asp(V²), D-Asp(V²) ou N-Me-Asp(V²) où V² est tel que défini ici, R¹⁶ a la même signification que celle définie pour R⁶ dans la revendication 1, Y¹ est U-Phe ou AcPhe où U est un groupe α -amino-protecteur, ou Y¹ est choisi dans le groupe suivant:

W¹-Val-Ile-Ser(V¹)-Asn-Ser(V¹)-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

W¹-Asn-Ser(V¹)-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

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W¹-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

AcThr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

AcAsn-Ser(V¹)-Phe,

W¹-Pro-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe,

W¹-Ser(V¹)-Pro-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe ou

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W¹-Ser(V¹)-Ser(V¹)-Pro-Thr(V¹)-Glu(V²)-Asn-Ser(V¹)-Phe où V¹ est un groupe protecteur pour le groupe hydroxyle de Thr ou Ser, V² est un groupe protecteur pour le ω -carboxyle de Glu et W¹ est un groupe α -amino-protecteur ou acétyle, et Z¹ est un groupe protecteur de carboxyle classique ou un support de résine; suivie par la déprotection (y compris le clivage du support de résine s'il est présent), et l'acylation et/ou l'amidification si nécessaire, du peptide protégé de formule 2 pour obtenir le peptide correspondant de formule 1; et, si on le souhaite, la transformation du peptide de formule 1 en un sel thérapeutiquement acceptable.

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8. Procédé pour préparer le peptide protégé de formule 2 selon la revendication 7 où Z¹ est un groupe protecteur de carboxyle ou un support de résine, qui comprend: le couplage par étapes dans l'ordre de la séquence d'acides aminés du peptide protégé de formule 2, des résidus d'acides aminés ou des fragments de peptide protégés dans lequel:

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i) les groupes latéraux labiles des résidus ou fragments sont protégés avec des groupes protecteurs appropriés pour empêcher des réactions chimiques de se produire à ce site jusqu'à ce que le groupe protecteur soit finalement retiré après la fin du couplage par étapes;

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ii) un groupe α -amino d'un corps réagissant qui se couple est protégé par un groupe α -amino-protecteur tandis que le groupe carboxyle libre de ce corps réagissant se couple avec le groupe α -amino libre du second corps réagissant; le groupe α -amino-protecteur étant un groupe qui peut être retiré sélectivement pour permettre à l'étape de couplage subséquente d'avoir lieu au niveau de ce groupe α -amino; et

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iii) le carboxyle C-terminal du résidu d'acide aminé ou fragment de peptide, qui doit devenir la fonction C-terminale du peptide protégé, est protégé avec un groupe protecteur approprié qui empêche une réaction chimique de se produire à ce site avant que la séquence d'acides aminés voulue pour le peptide protégé ait été assemblée;

pour obtenir le peptide protégé de formule 2.

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